Evaluation of Chlorpyrifos as a Toxic Air Contaminant

Consideration of the Addendum

Department of Pesticide Regulation
Human Health Assessment Branch

Scientific Review Panel Meeting
June 12, 2018
Presenting today:

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Today’s Presentation

Additional Data or Analysis added per request

Process for deriving a chlorpyrifos PoD for developmental neurotoxicity from in vivo animal data

Panel Discussion of Proposed Changes for Final Document

Review of TAC Authority

Document Scientific Sufficiency
Additional Data Included in Addendum
Additional Data included in Addendum

✅ Re-analysis of Registrant-submitted FIFRA Guideline Study (Hoberman, 1998)
  ▫ Special emphasis on brain morphology following in utero exposure
  ▫ Pages 6 – 12

✅ Thorough analysis of recent in vivo animal studies with developmental neurotoxicity outcomes
  ▫ Carr et al., 2017; Gómez-Giménez et al. (2017, 2018); Lee et al., 2015; Silva et al., 2017
  ▫ Pages 12 – 16 and 54 – 57

✅ Additional animal data
  ▫ Primate data (Coulston et al., 1971)
  ▫ Further genotoxic potential evaluation (Muller et al., 2014)
  ▫ Page 16 - 17
Additional Data included in Addendum, continued

✓ Most current epidemiological studies added
  ▪ Bulacan, Philippines; Central Ohio; Zhejiang Province, China; Mexico City
  ▪ Critical analysis of quantitative exposure analysis
  ▪ Pages 18 - 24

✓ New section on Delayed Neuropathy and Neurodegenerative Effects of Organophosphates added
  ▪ Includes human case reports or epidemiological studies, animal studies, and mechanistic studies for delayed neuropathy, Parkinson’s Disease, and Alzheimer’s Disease
  ▪ Page 24 - 47

✓ New section on additional human effects
  ▪ Includes respiratory effects (pages 48-53) and obesity (page 54)

✓ New section on recent advances in PBPK Modeling (page 55)
Additional Data included in Addendum, continued

- Added two additional age groups to the exposure assessment (infants and children 6 – 12 years old)
- New analysis of secondary drift exposure estimates
  - Based on DPR Air Monitoring Network data
  - Page 68
- Re-analysis of house dust data
  - Included new data from Gunier et al., 2016
  - Re-estimation on internal dose from 1999 house dust concentrations and 2006 house dust concentrations for comparison
  - Pages 68-70
- Revised Dietary Exposure Assessment
  - Updated risk values
  - Pages 71 - 74
Evaluation of Chlorpyrifos as a Toxic Air Contaminant

Addendum to the December 2017 Draft Evaluation
Hazard Identification
Hazard Identification - Introduction

PoD - The dose-response point that marks the beginning of a low-dose extrapolation; can be the lower bound on dose for an estimated incidence or a change in response level from a dose-response model (BMD), or a NOEL or LOEL for an observed incidence, or change in level of response.

Critical PoDs established from in vivo animal studies reporting DNT effects at dose levels lower than those that inhibit AChE.
Hazard Identification - Study Overviews

- Five recently published studies reported developmental toxicity in rodents (4 rat, 1 mice)
- Oral exposure route (three by gavage, two through the diet)
- Studies were not available to establish dermal and inhalation PoDs for DNT
- Two studies employed both gestational and lactational exposure
- Two studies employed direct pup exposure for either one or seven days starting at PND 10
- Neurodevelopmental responses in offspring were tested either in young pups (PNDs 21-25) or in adults (60-90 days)
Hazard Identification - Study Findings

• Three studies reported increased motor or total activity
• Two studies showed altered anxiety levels (decreased or increased)
• One study detected impaired spatial learning
• In four of studies, the LOEL was the lowest tested dose (0.1-0.5 mg/kg/day)
• Applying an uncertainty factor of 10 to those LOELs would result in an estimated no effect level (ENEL) for DNT of 0.01-0.05 mg/kg/day
• One study included a NOEL based on increased anxiety and motor activity in rats exposed in utero for 6 days (Silva et al., 2017)
Hazard Identification – Study Findings

- Only one study concurrently measured AChE activity
  - LOEL for brain AChE inhibition = 1.0 mg/kg/day (Carr et al., 2017)
- A registrant-submitted FIFRA Guideline Study (Hoberman, 1998)
  - Gestational and lactational exposure
  - RBC AChE inhibition was the most sensitive endpoint in this
    study with a BMDL10 / BMD10 of 0.03 / 0.06 mg/kg/day
  - HHA set the developmental LOEL at 1 mg/kg/day for brain
    morphometric changes in PND 66-71 females
  - This LOEL was 10 fold higher than the LOEL for DNT
    reported in the published studies
## Selected Animal Developmental Neurotoxicity Studies

<table>
<thead>
<tr>
<th>Species, Dosing Period, Doses (mg/kg/day)</th>
<th>Cholinesterase Inhibition</th>
<th>Developmental Neurotoxicity</th>
<th>Study</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Time tested</strong></td>
<td>LOEL</td>
<td>NOEL</td>
<td>Effects</td>
</tr>
<tr>
<td><strong>Species</strong></td>
<td>Plasma</td>
<td>RBC</td>
<td>Brain</td>
</tr>
<tr>
<td><strong>Gestation and postnatal exposure</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rat; Gavage GD 6-LD 11 0.3, 1.0, 5.0</td>
<td>Dam LD 22</td>
<td>0.3</td>
<td>--</td>
</tr>
<tr>
<td></td>
<td>Not tested</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Rat; Diet GD 7- PND 21 0.1, 0.3, 1.0</td>
<td>Not tested</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Rat; Diet GD 7- PND 21 0.1, 0.3, 1.0</td>
<td>Not tested</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td><strong>Gestation-only exposure</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rat; Gavage GD 14-20 0.01, 0.1, 1.0, 10</td>
<td>Not tested</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td><strong>Postnatal-only exposure</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rat; Gavage PND 10-16 0.5, 0.75 &amp; 1.0</td>
<td>Pups PND 16</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Mouse; Gavage PND 10 only 0.1, 1.0, 5.0</td>
<td>Pups PND 10</td>
<td>--</td>
<td>--</td>
</tr>
</tbody>
</table>
Hazard Identification - Point of Departure

- A NOEL of 0.01 mg/kg/day based on Silva et al., 2017 for increased anxiety and motor activity in rat pups
  - Supported by applying a 10-fold UF to the LOEL values (in the 4 other studies)
- Exposure duration in the 5 published studies varied from 1 to 35 days
  - Therefore, NOEL of 0.01 mg/kg/day could be applicable to acute and repeated exposures

\[ \therefore \text{ Acute oral PoD of } 0.01 \text{ mg/kg/day used to evaluate acute dermal and inhalation exposures using route-to-route extrapolation} \]
Exposure Assessment
Exposure Assessment

Comprised of two separate parts:

1. Bystander exposure to spray drift from chlorpyrifos applications

2. Dietary exposure to food and drinking water
Exposure Assessment - Bystander Drift

Calculating exposure estimates from:

1) Direct inhalation exposure from airborne chlorpyrifos resulting from pesticide application (1 hr exposure)

2) Incidental oral exposure (non-dietary hand-to-mouth, etc.) from chlorpyrifos that has deposited from spray drift on areas close to treated fields (1.5 hr exposure)

3) Dermal exposure through skin contact with contaminated soil/surfaces (1.5 hr exposure)

4) Combined spray drift exposure (dermal + non-dietary oral + inhalation)
Exposure Assessment - Bystander Drift, continued

We modeled:

- Distances of 25 – 2608 ft from edge of treated field
- Most common and reasonable “worst case” scenarios for application rates and volumes
- Aerial application (fixed wing and rotary)
- Aerial application modeled air concentrations were used as a surrogate for orchard airblast and ground boom
Exposure Assessment - Bystander Drift, continued

• Assessed four age groups
  ▪ Infant
  ▪ Child 1-2 years old
  ▪ Child 6-12 years old
  ▪ Females 13-49 years old

• Estimated absorbed doses for inhalation and dermal routes for MOE calculations
  ▪ For inhalation, assumed 100% external availability and 100% absorption
  ▪ For dermal, assumed 9.6% absorption
Exposure Assessment - Dietary

- Assessed four age groups
  - Infant
  - Child 1-2 years old
  - Child 6-12 years old
  - Females 13-49 years old

- Food (acute and steady state analyses)
  - From all foods with chlorpyrifos registrations
  - 79 individual tolerances and three crop group tolerances
  - Residues based on the USDA Pesticide Data Program monitoring database
  - Consumption based on 2003-2008 NHANES

- Drinking Water
  - Based on DPR measured residues in surface water in CA
Combined Exposure for 1 day

Spray Drift Exposure (dermal, non-dietary oral, inhalation)

Dietary Exposure (food + drinking water)

Spray Drift & Dietary Combined
## Margins of Exposure (MOE) from Exposure to Chlorpyrifos for Children 1-2yrs old

<table>
<thead>
<tr>
<th>Downwind Distance (ft)</th>
<th>Dermal</th>
<th>Incidental Oral</th>
<th>Inhalation</th>
<th>Spray Drift (1- 1.5 hr)</th>
<th>Dietary (1 day)</th>
<th>Combined Exposure</th>
<th>Spray Drift + Dietary</th>
</tr>
</thead>
<tbody>
<tr>
<td>25</td>
<td>2</td>
<td>8</td>
<td>9</td>
<td>1</td>
<td>24</td>
<td>54</td>
<td>1</td>
</tr>
<tr>
<td>50</td>
<td>2</td>
<td>10</td>
<td>10</td>
<td>2</td>
<td>24</td>
<td>54</td>
<td>2</td>
</tr>
<tr>
<td>100</td>
<td>3</td>
<td>15</td>
<td>13</td>
<td>3</td>
<td>24</td>
<td>54</td>
<td>2</td>
</tr>
<tr>
<td>250</td>
<td>7</td>
<td>29</td>
<td>19</td>
<td>5</td>
<td>24</td>
<td>54</td>
<td>4</td>
</tr>
<tr>
<td>500</td>
<td>12</td>
<td>54</td>
<td>30</td>
<td>9</td>
<td>24</td>
<td>54</td>
<td>6</td>
</tr>
<tr>
<td>1000</td>
<td>31</td>
<td>136</td>
<td>63</td>
<td>21</td>
<td>24</td>
<td>54</td>
<td>9</td>
</tr>
<tr>
<td>1320</td>
<td>52</td>
<td>232</td>
<td>92</td>
<td>33</td>
<td>24</td>
<td>54</td>
<td>11</td>
</tr>
<tr>
<td>2608</td>
<td>282</td>
<td>1255</td>
<td>279</td>
<td>140</td>
<td>24</td>
<td>54</td>
<td>15</td>
</tr>
</tbody>
</table>
Pesticide TAC Listing Criteria
For non-cancer effects, the threshold level is 10x below the air concentration which has been determined by the director to be protective of human health.

California Code of Regulations, Title 3, section 6864
Chlorpyrifos Evaluation as a TAC

TAC defined as air concentrations (modeled or monitored) that exceed the RfC/10

Chlorpyrifos will meet the criteria of listing as a TAC to protect against developmental neurotoxicity through both endpoints, DNT and AChE inhibition:

- TAC if air concentration > \(0.0005 \text{ mg/m}^3\) (500 ng/m\(^3\)) (Using DNT RfC for children 1-2 yrs of 0.005 mg/m\(^3\))

- TAC if air concentration > \(0.00095 \text{ mg/m}^3\) (950 ng/m\(^3\)) (Using AChEI RfC for children 1-2 yrs of 0.0095 mg/m\(^3\))
### Using Modeled Spray Drift Air Concentration (Addendum)

<table>
<thead>
<tr>
<th>Downwind Distance (ft)</th>
<th>1-hr TWA Concentration (mg/m³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>25</td>
<td>0.0493</td>
</tr>
<tr>
<td>50</td>
<td>0.0437</td>
</tr>
<tr>
<td>100</td>
<td>0.035</td>
</tr>
<tr>
<td>250</td>
<td>0.0237</td>
</tr>
<tr>
<td>500</td>
<td>0.0153</td>
</tr>
<tr>
<td>1000</td>
<td>0.0072</td>
</tr>
<tr>
<td>1320</td>
<td>0.00492</td>
</tr>
<tr>
<td>2608</td>
<td>0.00163</td>
</tr>
</tbody>
</table>

**Scenario:** Child 1-2 yrs with application by fixed wing aircraft with 2 gal/acre spray volume and 2 lbs/acre application rate

**Conclusion:** Modeled air concentrations at all distances exceeds the RfC/10 (i.e., > 0.0005 mg/m³ for DNT)
Using Measured Air Concentrations from AMN

**Conclusion:** Max 1-day measured air concentrations at Shafter in 2013 exceeded the RfC/10 in 2013 (i.e., > 0.0005 mg/m³ for DNT; > 500 ng/m³)

**Source:** DPR memo correlating agricultural use with ambient air concentrations of chlorpyrifos during 2011 – 2014 (Budahn, 2016)
Risk Characterization
Risk Characterization

• Risk for threshold effects is expressed as a margin of exposure (MOE)

• MOE = ratio of the critical NOEL or point of departure (PoD) to the estimated human exposure level

• Target MOE for developmental neurotoxic effects is 100
  ▪ 10x for interspecies sensitivity
  ▪ 10x for intraspecies variability

• Critical NOEL = 0.01 mg/kg/d
Route to route extrapolation

- Inhalation and dermal specific NOELs not available (no data to establish)

- Route-to-route extrapolation was performed to convert external dermal dose and inhalation concentrations to internal doses

- Derived acute inhalation and dermal PoDs from oral NOEL
Critical DNT NOELs used for the Risk Characterization of Chlorpyrifos (Table 23)

<table>
<thead>
<tr>
<th>Route</th>
<th>PoD&lt;sup&gt;a&lt;/sup&gt;</th>
<th>RfD&lt;sup&gt;b&lt;/sup&gt; or RfC</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Uncertainty Factors (UF)</strong></td>
<td></td>
<td>10 inter 10 intra 1 DNT</td>
</tr>
<tr>
<td><strong>Acute Oral [mg/kg/day]</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infants</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Children 1-2</td>
<td>0.01</td>
<td>0.0001</td>
</tr>
<tr>
<td>Children 6-12</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Females 13-49</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Acute Dermal [mg/kg/day]</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infants</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Children 1-2</td>
<td>0.104</td>
<td>0.001</td>
</tr>
<tr>
<td>Children 6-12</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Females 13-49</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Acute Inhalation [mg/m&lt;sup&gt;3&lt;/sup&gt;]</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Infants</td>
<td>0.405</td>
<td>0.004</td>
</tr>
<tr>
<td>Children 1-2</td>
<td>0.459</td>
<td>0.005</td>
</tr>
<tr>
<td>Children 6-12</td>
<td>0.624</td>
<td>0.006</td>
</tr>
<tr>
<td>Females 13-49</td>
<td>0.862</td>
<td>0.009</td>
</tr>
</tbody>
</table>
Margins of Exposure

- Combined spray drift exposures estimates at 2608 feet for dermal, incidental oral, and inhalation routes were combined with the 99.9\textsuperscript{th} percentile exposures from dietary and drinking water for chlorpyrifos.

- At 2608 feet from a field treated with chlorpyrifos, the combined spray drift MOEs for all four sensitive population subgroups were at or greater than the target of 100.

- However, when dietary and drinking water exposures were added, the aggregate MOEs for these combined routes and sources of exposure were below all the target of 100 (indicating a health concern).
Margins of Exposure using DNT NOEL for Combined Spray Drift, Dietary & Drinking Water (2608ft from Field Edge)

<table>
<thead>
<tr>
<th>Population Subgroup</th>
<th>Margin of Exposure&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Diet Only&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>All Infants &lt; 1 year</td>
<td>37</td>
</tr>
<tr>
<td>Children 1-2 years</td>
<td>24</td>
</tr>
<tr>
<td>Children 6-12 years</td>
<td>53</td>
</tr>
<tr>
<td>Females 13-49 years</td>
<td>67</td>
</tr>
</tbody>
</table>

Abbreviations: DNT = Developmental Neurotoxicity, NOEL = No-Observed-Effect Level.

<sup>a</sup> Margin of Exposure (MOE) = NOEL / Exposure; DNT NOEL = 0.01 mg/kg/day based on changes in cognition, motor control and behavior in rats and mice (Lee et al, 2015, Silva et al, 2017, Carr et al, 2017, Gómez-Giménez, 2017, 2018).

<sup>b</sup> Dietary exposure estimate at the 99.9<sup>th</sup> percentile was used in the MOE calculation.

<sup>c</sup> Drinking water exposure estimate based on the 99.9<sup>th</sup> percentile from DPR surface water monitoring was used in the MOE calculation.

<sup>d</sup> Combined Spray Drift MOE is the MOE for the combined dermal, incidental oral and inhalation exposure from spray drift at 2608 ft from the treated field which is the only distance where MOEs were greater than 100 for all routes (see Table 24).

<sup>e</sup> Combined MOE = DNT NOEL (0.01) / (Diet + Drinking Water + Combined Spray Drift) Exposure. Red shading indicates MOEs that are below the target of 100.
Proposed additions or edits for final TAC Evaluation Document
For Exposure and TAC determination discussion in Addendum

1. Clearly state how chlorpyrifos can meet the TAC criteria
   - Addition in Risk Characterization, Risk Appraisal, and Conclusion sections
   - Option: reserve discussion on TAC designation for AChE inhibition endpoint in appendix

2. Clearly state how inhalation RfC using DNT endpoint meets TAC criteria — however — consumption of food and drinking water drive the risk
   - Addition in Risk Characterization, Risk Appraisal, and Conclusion sections
For Discussion of DNT Endpoint in Addendum

1. Move comparison of PoDs and reference concentrations/ doses of developmental neurotoxic effect vs. AChE inhibition to front matter of Appendix 3

2. Revise Executive Summary and Conclusion to focus on DNT
   - Comprehensive analysis of all currently available data to establish a PoD based directly on developmental neurotoxicity
Scientific Sufficiency of the TAC Evaluation
Scientific Review Panel on Toxic Air Contaminants

The Panel shall review “the scientific data on which the report is based, the scientific procedures and methods used to support the data, and the conclusions and assessments on which the report is based.”

If SRP determines “the health effects report is seriously deficient,” it returns the report to the DPR director who shall revise and resubmit it within 30 days of receiving the SRP’s determination of deficiency, and prior to developing control measures or other regulations.

Food and Agricultural Code Section 14023(b)-(c)
Scientific Sufficiency of the TAC Evaluation

• The report shall assess (FAC § 14023(a)):
  ▪ Availability and quality of data on health effects
  ▪ Potency, mode of action, and other relevant biological factors
  ▪ An estimate of the levels of exposure that may cause or contribute to adverse health effects
  ▪ The range of risk to humans resulting from current or anticipated exposure.
Thank you